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- (71) Applicant: **NANOPHASE TECHNOLOGIES CORPORATION** [US/US]; 1319 Marquette Drive, Romeoville, IL 60446 (US).
- (72) Inventors: **CLAYTON, Roger, H.**; 856 South Wright Street, Naperville, IL 60540 (US). **BROTZMAN, Richard, W., Jr.**; 318 Kent Court, Naperville, IL 60540-5711 (US). **MURRAY, Patrick, G.**; 1737 Columbine Court, Yorkville, IL 60560 (US).
- (74) Agent: **RUPERT, Douglas, S.**; Wildman, Harrold, Allen & Dixon, 225 West Wacker Street, Suite 2800, Chicago, IL 60606 (US).

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(54) Title: STABLE DISPERSIONS OF NANOPARTICLES IN AQUEOUS MEDIA

(57) Abstract: A process to prepare a stable dispersion of nanoparticles in aqueous media. A dispersant and aqueous are combined to form a mixture. The dispersant is selected from the group comprising copolymers and cyclic phosphates. Nanoparticles are added to the mixture to form the dispersion.

STABLE DISPERSIONS OF NANOPARTICLES IN AQUEOUS MEDIA

FIELD OF THE INVENTION

5 The present invention relates to dispersions of nanoparticles in aqueous media, and more specifically to stable aqueous dispersions of nanocrystalline metals and metal oxides.

BACKGROUND OF THE INVENTION

 Stable aqueous-based dispersions of nanoparticles, such as substantially
10 spherical nanocrystalline metals and/or metal oxides would be useful for many applications. Such dispersions could serve as a component of transparent coatings, which could be used on surfaces to yield unique properties such as abrasion resistance, radiation absorption or reflection, electrical conductivity, and catalytic function. Other applications of dispersions include, but are not limited to, functioning as abrasive or
15 polishing fluids, thermal transfer fluids, catalytic additives, ingredients to cosmetic and personal care formulations, and electro-rheological fluids.

 Generally products utilizing the dispersions described above have different pH values than the natural pH of metal and/or metal oxides in water. This often leads to dispersion instability because, as the dispersion pH is adjusted for application use, the
20 isoelectric point of the dispersed phase is encountered and flocculation of the nanoparticles is initiated. Thus, it would be desirable to form stable aqueous-based dispersions at pH values required by the application, especially pH values above or near the isoelectric point of the metal and/or metal oxide. Therefore, a need exists for a method of preparation of stable dispersions of nanoparticles, such as substantially

spherical nanocrystalline metals and/or metal oxides, and aqueous media at a variety of pH values.

SUMMARY OF THE INVENTION

5 In one example, the present invention relates to a method of preparing or forming stable dispersions of nanoparticles and aqueous media. The method comprises combining a dispersant with aqueous media to form a mixture. The dispersant in one example is selected from the group comprising water soluble copolymers and cyclic phosphates. Nanoparticles, such as substantially spherical nanocrystalline metal and/or
10 metal oxide particles are added to the mixture.

DETAILED DESCRIPTION OF THE INVENTION

Following are definitions of terms that are used throughout the description:

Isoelectric point – the pH of zero net charge on a nanoparticle in dispersion. The isoelectric point is determined by measuring the zeta-potential of a nanoparticle
15 dispersion and a buffer to maintain dispersion pH. The pH where the zeta-potential is zero is the isoelectric point.

Long-term stable dispersion – the dispersed nanoparticles do not aggregate (no increase in particle size) and gravitational sedimentation is minimized on the time frame of 6 months and longer.

20 Short-term stable dispersion – the dispersed nanoparticles are initially well dispersed but begin to aggregate, displaying an increased particle size and concomitant sedimentation, on the time frame of days to weeks.

Water-soluble dispersants are used in a method of dispersing nanoparticles, such as substantially spherical metal and/or metal oxide nanoparticles. In one example,

the nanoparticles comprise the nanocrystalline materials described in U.S. Patent Number 5,874,684, entitled "Nanocrystalline Materials", which was granted to Parker et al. on February 23, 1999, and which is hereby incorporated by reference. The aqueous-based dispersions, of the present invention, are made by dissolving dispersant
5 in water and adding the nanoparticles while high shear mixing (e.g., ultrasonication, rotor-stator mixing, homogenizer mixing, etc.) Substantially spherical nanocrystalline metals and/or metal oxides are dispersed above their isoelectric points using a variety of water soluble dispersants, including but not limited to, pigment dispersants, surfactants, wetting agents, coupling agents (hereinafter referred to collectively in this
10 document as "dispersants"). The dispersants range from small molecules to oligomeric materials to polymers to coupling agents and featured a variety of different surface anchoring groups (acidic, basic, or neutral), and had different ionic character (cationic, anionic, or neutral).

Screenings were conducted utilizing the dispersants to disperse substantially
15 spherical nanocrystalline metals and metal oxides. Experiments were constructed to cover a number of different particle concentrations as well as a number of different dispersant levels with respect to the particle. Samples were prepared by ultrasonication and the quality of dispersion was measured by the following criteria:

1. Qualitative appearance of the dispersion
- 20 2. Particle size determination
3. Dispersion stability with respect to gravimetric sedimentation over time

Surfactants, such as those given in the examples which follow, were employed to obtain stable dispersions of substantially spherical nanocrystalline metal and metal

oxide particles. The pH was adjusted above the isoelectric point of the particles with hydroxide bases. Surprisingly, only water-soluble copolymers and, for some nanoparticles, cyclic phosphates, were found to yield stable aqueous-based dispersions of substantially spherical nanocrystalline metals and/or metal oxides above the
5 isoelectric point of the particles. The resulting aqueous-based dispersions of substantially spherical nanocrystalline particles are stable, have a pH greater than the isoelectric point of the particles in an aqueous-based medium, and could be incorporated into application formulations without inducing flocculation of the particles.

10 A description of several exemplary experiments now follows for illustrative purposes.

**Example 1: Aqueous-Based Dispersions of Substantially Spherical
Nanocrystalline Aluminum Oxide**

15 Dispersants evaluated in aqueous-based dispersions of aluminum oxide are listed in Table 1. Commercial dispersant names, maximum weight percent oxide in a fluid dispersion, weight percent dispersant with respect to aluminum oxide, mean particle size in dispersion on a volume-weight basis in dispersions as made, dispersion stability after the dispersion pH was increased above the isoelectric point of aluminum
20 oxide dispersion using hydroxide bases (stable dispersion = S, long term – LT, short term – ST, flocculated dispersion = F), and dispersant type are tabulated. The dispersions that were initially stable were monitored over time and were further characterized. The general dispersion effectiveness falls into two groups depending on the length of time the dispersion remains stable. Long-term stable dispersions are
25 stable for at least 6 months and do not exhibit aggregation and particle size growth.

However, short-term stable dispersions exhibit aggregation and particle size growth on the time frame of days to weeks.

- Only water-soluble copolymers that have polymer segments that are attractive to the nanocrystalline particle and different polymer segments that render them water-soluble yield long-term stable dispersions. This is a surprising result – homopolymers of acrylic acid as a class only render the dispersions stable for short times.

Table 1. Dispersants for Aqueous-Based Aluminum Oxide Dispersions					
Dispersant	Max Oxide wt%	Disp wt%	Initial PS, nm <d>vol	Dispersion Stability	Dispersant Type
Long Term Stable					
Polyacryl C50-45AN	65	10	135	S – LT	Acrylamidomethylpropane sulfonic acid / acrylic acid copolymer, neutral to pH = 8
Tego 752W	65	10	135	S – LT	Maleic acid/vinyl polyether copolymer, pH = 6
Disperbyk-190	50	10	135	S – LT	Non-Ionic copolymer with carboxy anchor groups, pH= 7
Zephrym PD3315	50	10	135	S – LT	Propylene oxide/acrylic acid copolymer, pH = 8
Short Term Stable					
Hydropalat 44	20	10	150	S – ST	Acrylic acid homopolymer, pH = 7.8
Polacryl A60-40S	20	10	150	S – ST	Acrylic acid homopolymer, pH = 8.0
Polacryl B55-50AN	20	10	150	S – ST	Acrylic acid homopolymer, pH = 6.5
Polacryl A60-35S	40	10	150	S – ST	Acrylic acid homopolymer, pH = 8.0
Hydropalat 100	10	20	150	S – ST	Acrylic acid homopolymer, pH = 6.5
HMP	20	2	150	S – ST	Sodium hexametaphosphate, ring structure
Unstable					
Solsperse 27000	0	5	> 500	F	Nonionic polymer
PVP K-15	0	10	> 500	F	Polyvinylpyrrolidone, MW = 9700
Ganex P-904 LC	0	5	> 500	F	90% PVP/10% Poly-C4, HLB = 18-20
Solsperse 20000	0	10	> 500	F	Cationic polymer
Solsperse 40000	0	20	> 500	F	Anionic polymer neutralized with DEA
Solsperse 41090	0	20	> 500	F	Anionic polymer, pH = 2 – 3
PVP/VA S-630	0	10	> 500	F	Polyvinylpyrrolidone/Vinyl acetate
Hydropalat 3216	0	20	> 500	F	Nonionic and Ionic Surfactants

Example 2: Aqueous-Based Dispersions of Substantially Spherical Nanocrystalline Cerium Oxide

Dispersants evaluated in aqueous-based dispersions of cerium oxide are listed in Table 2. Commercial dispersant names, weight percent oxide in dispersion, weight percent dispersant with respect to cerium oxide, mean particle size in dispersion on a volume-weight basis in dispersions as made, dispersion stability after the dispersion pH was increased above the isoelectric point of cerium oxide dispersion using hydroxide bases (stable dispersion = S, long term – LT, short term – ST, flocculated dispersion = F), and dispersant type are tabulated. The dispersions that were initially stable were evaluated over time and were further characterized. As with alumina, the general dispersion effectiveness for ceria falls into two groups depending on the length of time the dispersion remains stable – long-term and short-term stable dispersions.

Only water-soluble copolymers that have polymer segments that are attractive to the nanocrystalline particle and polymer segments that render them water-soluble yield long-term stable dispersions. This is a surprising result – homopolymers of acrylic acid as a class only render the dispersions stable for short times. In the case of unstable dispersions the observed flocculation is irreversible.

Table 2. Dispersants for Aqueous-Based Cerium Oxide Dispersions					
Dispersant	Oxide wt%	Disp wt%	Initial PS, nm <d>vol	Dispersion Stability	Dispersant Type
Long-Term Stable					
Polyacryl C50-45AN	20	10	120	S – LT	Acrylamidomethylpropane sulfonic acid / acrylic acid copolymer, neutral to pH = 8
Tego 752W	20	10	120	S – LT	Maleic acid/vinyl polyether copolymer, pH = 6
Disperbyk-190	20	10	120	S – LT	Non-Ionic copolymer with carboxy anchor groups, pH= 7
Zephrym PD3315	20	10	120	S – LT	Propylene oxide/acrylic acid copolymer, pH = 8
Short-Term Stable					

Polacryl A60-35S	20	10	150	S – ST	Acrylic acid homopolymer, pH = 8.0
Polacryl A60-40S	20	10	150	S – ST	Acrylic acid homopolymer, pH = 8.0
Polacryl B55-50AN	20	10	150	S – ST	Acrylic acid homopolymer, pH = 6.5
Polacryl B55-50A	20	10	150	S – ST	Acrylic acid homopolymer
Hydroplat 100	20	10	150	S – ST	Hydrophobically modified acrylic acid homopolymer
Unstable					
PVP K-15	0	20	> 500	F	Polyvinylpyrrolidone, MW = 9700
Solsperse 27000	0	10	> 500	F	Nonionic polymer
PVP/VA S-630	0	10	> 500	F	Polyvinylpyrrolidone/Vinyl acetate
Ganex P-904 LC	0	5	> 500	F	90% PVP/10% Poly-C4, HLB = 18-20
HMP	0	2	> 500	F	Sodium hexametaphosphate, ring structure
Solsperse 40000	0	20	> 500	F	Anionic polymer neutralized with DEA
Solsperse 41090	0	20	> 500	F	Anionic polymer, pH = 2 - 3

Example 3: Aqueous-Based Dispersions of Substantially Spherical Nanocrystalline Zinc Oxide

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Dispersants evaluated in aqueous-based dispersions of zinc oxide are listed in Table 3. Commercial dispersant names, maximum weight percent oxide in fluid dispersion, weight percent dispersant with respect to zinc oxide, mean particle size in dispersion on a volume-weight basis in dispersions as made, dispersion stability after the dispersion pH was increased above the isoelectric point of zinc oxide using hydroxide bases (stable dispersion = S, long term – LT, short term – ST, flocculated dispersion = F), and dispersant type are tabulated. The dispersions that were initially stable were evaluated over time and were further characterized. As with alumina and ceria, the general dispersion effectiveness for ceria falls into two groups depending on the length of time the dispersion remains stable – long-term and short-term stable dispersions.

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Only water-soluble copolymers that have polymer segments that are attractive to the nanocrystalline particle and polymer segments that render them water-soluble

yield long-term stable dispersions. This is a surprising result – homopolymers of acrylic acid as a class only render the dispersions stable for short times.

Table 3. Dispersants for Aqueous-Based Zinc Oxide Dispersions

Dispersant	Max Oxide, wt%	Disp, wt%	PS, nm <d>vol	Dispersion Stability	Dispersant Type
Long-Term Stable					
Polyacryl C50-45AN	40	5	310	S - LT	Acrylamidomethylpropane sulfonic acid / acrylic acid copolymer, neutral to pH = 8
Disperbyk 190	60	4	310	S - LT	Non-Ionic copolymer with carboxy anchor groups, pH= 7
Zephrym PD3315	40	3	310	S - LT	Propylene oxide / acrylic acid copolymer, pH = 8
HMP	30	2	310	S - LT	Sodium hexametaphosphate, ring structure
Short-Term Stable					
Zephrym PD 3076	28	10	250	S - ST	Acrylic acid-based homopolymer
Hydropalat 44	30	0.7	390	S - ST	Acrylic acid homopolymer, pH = 7.8
Hydropalat 100	30	20	430	S - ST	Acrylic acid homopolymer, pH = 6.5
Polacryl A60-35S	40	5	390	S - ST	Acrylic acid homopolymer, pH = 8.0
Polacryl A60-40S	40	5	370	S - ST	Acrylic acid homopolymer, pH = 8.0
Polacryl B55-50AN	40	5	300	S - ST	Acrylic acid homopolymer, pH = 6.5
Unstable					
Polacryl B55-50A	0	10	> 500	F	Acrylic acid homopolymer, pH = 3.5
PVP K-15	0	10	> 500	F	Polyvinylpyrrolidone, MW = 9700
Hydropalat 3216	0	20	> 500	F	Nonionic and Ionic Surfactants
Solsperse 20000	0	5	> 500	F	Basic, cationic single anchor, single polymer chain
Solsperse 27000	0	5	> 500	F	Nonionic polymer
Solsperse 40000	0	20	> 500	F	Anionic polymer neutralized with DEA
Solsperse 41090	0	18	> 500	F	Anionic polymer
Zephrym PD 3800	0	10	> 500	F	Nonionic/Anionic Surfactant blend
Zephrym PD 3100	0	10	> 500	F	Alcohol ethoxylate
Zephrym PD 7000	0	10	> 500	F	Nonionic surfactant
Zephrym PD 2434	0	20	> 500	F	Polymeric dispersant
Disperbyk 184	0	20	> 500	F	Polymeric dispersant
Disperbyk 192	0	20	> 500	F	Polymeric dispersant
PVP/VA S-630	0	10	> 500	F	Polyvinylpyrrolidone/Vinyl acetate
Ganex P-904 LC	0	10	> 500	F	90% PVP/10% Poly-C4, HLB = 18-20
Copolymer 958	0	11	> 500	F	PVP/Dimethylaminoethylmethacrylate copolymer
PVP/VA W-635	0	10	> 500	F	PVP/vinyl acetate copolymer
Hydropalat 188A	0	20	> 500	F	Polyethyleneglycol dioleate, Nonionic

					surfactant
Hydropalat 535N	0	20	> 500	F	Oleoalkylenoxide block copolymer
Hydropalat 1080	0	20	> 500	F	Oleoalkylenoxide block copolymer
Zonyl FSO100	0	2	> 500	F	Fluorinated surfactant
Alkox E-30	0	10	> 500	F	Polyethyleneoxide
Alkox E-160	0	10	> 500	F	Polyethyleneoxide
Alkox R-150	0	20	> 500	F	Polyethyleneoxide
Alkox R-400	0	20	> 500	F	Polyethyleneoxide

**Example 4: Aqueous-Based Dispersions of Other Substantially Spherical
Nanocrystalline Particles – Copper Oxide, Silver, Antimony Tin
Oxide, Indium Tin Oxide**

5 Long-term stable, aqueous-based dispersions of other substantially spherical nanocrystalline particles – copper oxide, silver, antimony tin oxide, indium tin oxide – are produced using water-soluble copolymer dispersant levels from 1 to 20-wt% dispersant with respect to nanocrystalline particles, depending on the copolymer dispersant used. The copolymer dispersant stabilizes the volume-weighted mean
10 particle size preventing aggregation (the formation of grape-like clusters).

**Example 5: The Stability of Aqueous-Based Dispersions of Substantially
Spherical Nanocrystalline Cerium Oxide**

15 The mean particle size, of substantially spherical ceria, in aqueous dispersion at pH 7.5 on a volume-weight basis (measured using dynamic light scattering), as functions of time and dispersant type, are given in Table 4. The mean particle size is stable for Zephrym PD 3315 and Polyacryl C50-45AN, water-soluble copolymers that have polymer segments that are attractive to the nanocrystalline particle and polymer
20 segments that render them water-soluble. Where as the mean particle size grows over time for Polyacryl B55-50AN and Hydropat 44, homopolymers of acrylic acid. This

is a surprising result. – homopolymers of acrylic acid as a class are claimed to render the dispersions stable (see US Patent 5,876,490)

Dispersant	PS 0 days	PS 1 day	PS 3 days	PS 21 days	PS 54 days	PS 12 mo
Hydropatat 44	289 nm	268 nm	330 nm	327 nm	402 nm	378 nm
Polacryl B55-50AN	155 nm	171 nm	152 nm	315 nm	376 nm	415 nm
Zephrym PD 3315	173 nm	212 nm	141 nm	163 nm	200 nm	216 nm
Polyacryl C50-45AN	178 nm	155 nm	146 nm	172 nm	180 nm	196 nm

5

Example 6. Settling Stability of Aqueous Dispersions of Substantially Spherical Nanocrystalline Ceria at Elevated pH

The stability of aqueous dispersions of substantially spherical nanocrystalline ceria at elevated pH with respect to gravitational sedimentation was quantified as a function of dispersant type, dispersant concentration, and pH. A slow rate of gravitational sedimentation is desired in storage containers to minimize the amount of mixing required to homogenize the concentration. For aqueous ceria dispersions the problem is particularly challenging since the density of the ceria is approximately seven times the density of water and for 20-wt% ceria dispersions the dispersion viscosity is less than 10 cP.

Dispersions were prepared using C50-45AN and B55-50AN. Each sample in Table 5 was placed into a 500 mL polypropylene graduated cylinder. The cylinder contained a column of ceria dispersion 27.5 cm high. The graduated cylinder was covered tightly with Parafilm and set aside for 30 days.

20

Table 5. Dispersion Samples in Gravimetric Sedimentation Study		
Sample	weight % Dispersant (based on ceria)	initial d<vol> nm (SD nm)
114A	8 % C50-45AN	94 (21)
114B	9 % C50-45AN	93 (21)
114C	10 % C50-45AN	94 (21)
114D	11 % C50-45AN	92 (20)
114E	12 % C50-45AN	94 (21)
114F	10 % B55-50AN	94 (22)

* Horiba LA-910; mean volume weighted PS and standard deviation

5

After thirty days, 100 mL aliquots (5.5 cm of dispersion) of the ceria dispersion were carefully removed from the cylinder. These aliquots were taken from the top of the cylinder with a polypropylene syringe equipped with a virgin 6" stainless steel needle, located just beneath the surface of the liquid in a fashion such that the liquid below was not disturbed. Each 100 mL aliquot was stored in a separate 125 mL polypropylene container and named "1" through "5" depending on where in the graduated cylinder it was taken. For example, 114A-1 was taken from the top of the graduated cylinder while 114A-5 was taken from the bottom of the graduated cylinder. Each 100 mL aliquot was characterized by the loss on drying and by Horiba particle size determination. The amount of sediment that would not pour out of the graduated

cylinder after 20 seconds of inversion was also determined. These data are presented in Table 6.

Table 6. Sediment, solids, and PS for Table 5 Dispersions			
Sample	sediment (g)	% solids (LOD)	d<vol>, nm (SD, nm)*
114A-1		10.1	85 (16)
114A-2		15.2	95 (20)
114A-3		16.6	103 (22)
114A-4		17.0	105 (23)
114A-5		21.2	108 (25)
114A-sediment	10.04	-	-
114B-1		9.8	86 (16)
114B-2		15.2	96 (20)
114B-3		16.4	103 (23)
114B-4		16.7	105 (24)
114B-5		20.2	108 (25)
114B-sediment	9.06	-	
114C-1		10.9	86 (17)
114C-2		15.6	97 (20)
114C-3		16.3	104 (23)
114C-4		17.1	106 (24)
114C-5		21.4	109 (25)
114C-sediment	6.78	-	
114D-1		10.5	86 (16)
114D-2		15.8	96 (20)
114D-3		16.7	103 (22)
114D-4		16.9	106 (24)
114D-5		20.7	108 (25)
114D-sediment	6.94	-	

114E-1		11.5	86 (17)
114E-2		16.1	98 (21)
114E-3		17.0	105 (23)
114E-4		17.2	106 (24)
114E-5		21.2	111 (27)
114E-sediment	7.06	-	
114F-1		7.5	84 (16)
114F-2		9.3	87 (17)
114F-3		9.5	88 (17)
114F-4		9.3	89 (18)
114F-5		21.6	120 (37)
114F-sediment	51.5	-	

* Horiba LA-910; mean volume weighted PS and standard deviation

- 5 Data in Table 6 show the amount of sediment in C50-45AN samples decreases until 10% C50-45AN is reached, after which there is little improvement to be gained by adding more dispersant. The sediment obtained with the dispersant B55-50AN, a homopolymer of acrylic acid, at 10% by weight (51.5%) is by far greater than C50-45AN at any concentration examined.
- 10 Although various examples have been depicted and described in detail herein, it will be apparent to those skilled in the relevant art that various modifications, additions, substitutions and the like can be made without departing from the spirit of the invention and these are therefore considered to be within the scope of the invention defined.

Claims

What is claimed is:

1. A process to prepare a stable dispersion of nanoparticles in aqueous media, the
5 process comprising:
combining a dispersant with the aqueous media to form a mixture, wherein
the dispersant is selected from the group comprising copolymers and cyclic
phosphates; and
adding nanoparticles to the mixture.
- 10 2. The process of claim 1, further comprising:
selecting one of metal oxides and mixed metal oxides as the nanoparticles to
add to the mixture.
3. The process of claim 2, further comprising:
selecting metal oxides from a group comprising aluminum oxide, zinc
15 oxide, iron oxide, cerium oxide, chromium oxide, antimony tin oxide, and
indium tin oxide as the nanoparticles to add to the mixture.
4. The process of claim 1, further comprising:
selecting one of substantially spherical nanocrystalline metal oxides and
substantially spherical nanocrystalline mixed metal oxides as the
20 nanoparticles to add to the mixture.
5. The process of claim 1, further comprising:
selecting the nanoparticles generally to have a size distribution and range in
mean diameter from about 1 nm to about 900 nm.
6. The process of claim 5, wherein the selecting step comprises:

selecting the nanoparticles generally to have a size distribution and range in mean diameter from about 2 nm to about 100 nm.

7. The process of claim 6, wherein the selecting step comprises:

selecting the nanoparticles generally to have a size distribution and range in mean diameter from about 5 nm to about 40 nm.

8. The process of claim 1, further comprising:

selecting the dispersant to be a copolymer having one or more functional groups capable of anchoring to a surface of at least one of the nanoparticles.

9. The process of claim 8, wherein the dispersant anchors to the nanoparticle surface through at least one of acidic interactions, basic interactions, neutral interactions, and covalent interactions.

10. The process of claim 9, wherein interaction between the dispersant and the at least one of the nanoparticles is of one of cationic character, anionic character, and neutral character.

11. The process of claim 1, wherein the dispersant is soluble in the aqueous media.

12. The process of claim 1, wherein the dispersant is a cyclic phosphate.

13. The process of claim 1, wherein the step of combining comprises:

mixing the dispersant to the aqueous media.

14. The process of claim 13 wherein the step of mixing is accomplished through one of high-shear mixing and ultrasonic mixing of the dispersant to the aqueous media.

15. The process of claim 1, wherein the step of adding comprises:

mixing the nanoparticles with the mixture.

16. The process of claim 15, wherein the step of adding is accomplished through one of high-shear mixing and ultra-sonic mixing the nanoparticles with the mixture.
17. A composition of nanoparticles dispersed in aqueous media produced by the
5 process of claim 1.
18. The composition of claim 17, further comprising:
selecting one of metal oxides and mixed metal oxides as the nanoparticles.
19. The composition of claim 18, further comprising:
selecting metal oxides from a group comprising aluminum oxide, zinc
10 oxide, iron oxide, cerium oxide, chromium oxide, antimony tin oxide, and indium tin oxide as the nanoparticles to add to the mixture.
20. The composition of claim 17, further comprising:
selecting one of substantially spherical nanocrystalline metal oxides and substantially spherical nanocrystalline mixed metal oxides as the
15 nanoparticles to add to the mixture.
21. The composition of claim 17, further comprising:
selecting the nanoparticles generally to have a size distribution and range in mean diameter from about 1 nm to about 900 nm.
22. The composition of claim 21, wherein the selecting step comprises:
20 selecting the nanoparticles generally to have a size distribution and range in mean diameter from about 2 nm to about 100 nm.
23. The composition of claim 22, wherein the selecting step comprises:
selecting the nanoparticles generally to have a size distribution and range in mean diameter from about 5 nm to about 40 nm.

24. The composition of claim 17, further comprising:
selecting the dispersant to be a copolymer.
25. The composition of claim 24, further comprising:
selecting the dispersant to have one or more functional groups capable of
5 anchoring to a surface of at least one of the nanoparticles.
26. The composition of claim 25, wherein the copolymeric dispersant anchors to the
nanoparticle surface through at least one of acidic interactions, basic
interactions, neutral interactions, and covalent interactions.
27. The composition of claim 26, wherein interaction between the copolymeric
10 dispersant and the at least one of the nanoparticles is of one of cationic
character, anionic character, and neutral character.
28. The composition of claim 17, wherein the dispersant is soluble in the aqueous
media.
29. The composition of claim 17, wherein the dispersant is cyclic phosphate-based.